



Aerosol properties and their influences on low warm clouds during the Two-Column Aerosol Project

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Abstract. Twelve months of measurements collected during the Two-Column Aerosol Project field campaign at Cape Cod, Massachusetts, which started in the summer of 2012, were used to investigate aerosol physical, optical, and chemical properties and their influences on the dependence of cloud development on thermodynamic (i.e., lower tropospheric stability, LTS) conditions. Relationships between aerosol loading and cloud properties under different dominant air-mass conditions and the magnitude of the first indirect effect (FIE), as well as the sensitivity of the FIE to different aerosol compositions, are examined. The seasonal variation in aerosol number concentration (N_a) was not consistent with variations in aerosol optical properties (i.e., scattering coefficient, σ_s , and columnar aerosol optical depth). Organics were found to have a large contribution to small particle sizes. This contribution decreased during the particle growth period. Under low-aerosol-loading conditions, the liquid water path (LWP) and droplet effective radius (DER) significantly increased with increasing LTS, but, under high-aerosol-loading conditions, LWP and DER changed little, indicating that aerosols significantly weakened the dependence of cloud development on LTS. The reduction in LWP and DER from low- to high-aerosol-loading conditions was greater in stable environments, suggesting that clouds under stable conditions are more susceptible to aerosol perturbations than those under more unstable conditions. High aerosol loading weakened the increase in DER as LWP increased and strengthened the increase in cloud optical depth (COD) with increasing LWP, resulting in changes in the interdependence of cloud properties. Under both continental and marine air-mass conditions, high aerosol loading can

significantly increase COD and decrease LWP and DER, narrowing their distributions. Magnitudes of the FIE estimated under continental air-mass conditions ranged from 0.07 ± 0.03 to 0.26 ± 0.09 with a mean value of 0.16 ± 0.03 and showed an increasing trend as LWP increased. The calculated FIE values for aerosols with a low fraction of organics are greater than those for aerosols with a high fraction of organics. This implies that clouds over regions dominated by aerosol particles containing mostly inorganics are more susceptible to aerosol perturbations, resulting in larger climate forcing, than clouds over regions dominated by organic aerosol particles.

1 Introduction

Aerosols can significantly influence climate change through their direct and indirect effects (IPCC, 2013; Li et al., 2016, 2017). The aerosol direct effect is when aerosol particles change Earth's radiative balance by scattering and absorbing solar radiation (e.g., Liu et al., 2012; Yang et al., 2016, 2018). The aerosol indirect effect is when aerosols change cloud microphysical, macrophysical, and precipitation properties through their role as cloud condensation nuclei (CCN) or ice nuclei (IN). Under constant liquid water path (LWP) conditions, an increase in aerosol concentration will lead to an increase in CCN concentration. This results in an increase in cloud droplet number concentration, a decrease in the cloud droplet effective radius (DER), and a more reflective cloud. This is referred to as the first aerosol indirect effect (FIE) (Twomey, 1977; Feingold et al., 2003; Garrett et

al., 2004; Zhao et al., 2018, 2019). The decrease in DER will reduce the chances of precipitation forming, which prolongs the lifetime of a cloud and enhances its LWP. This is known as the second aerosol indirect effect (e.g., Albrecht, 1989). Aerosols also influence cloud properties through the thermal emissivity effect (e.g., Garrett and Zhao, 2006; Zhao and Garrett, 2015) and the semidirect effect (e.g., Koren et al., 2004). Estimates of indirect aerosol effects have large uncertainties (Boney and Dufresne, 2006; Lohmann et al., 2010). This makes the impact of aerosols on the prediction of the current and future behaviors of Earth's climate system highly uncertain (McComiskey and Feingold, 2008; IPCC, 2013).

The observed response of warm low cloud properties to aerosol properties has been observed from satellite-based remote sensing (Bréon et al., 2002; Lebsack et al., 2008; Su et al., 2010; Wang et al., 2014), surface-based remote sensing (Kim et al., 2003; Feingold et al., 2003, 2006; Garrett et al., 2004; McComiskey et al., 2009; Liu et al., 2016; Qiu et al., 2017; Liu and Li, 2018a), combined surface measurements and satellite retrievals (Sporre et al., 2012, 2014), and aircraft measurements (Zhang et al., 2011; Twohy et al., 2013; Painemal and Zuidema, 2013; Werner et al., 2014; Zhao et al., 2018, 2019; Yang et al., 2019). Most of these studies have shown that DER significantly decreases as aerosol loading increases. However, LWP can increase or decrease with aerosol loading, depending on cloud thermodynamics and dynamics (Han et al., 2002). Current estimates of FIE from all available observational platforms have a large range of values because each set of measurements used has its own set of uncertainties, and so do their approaches. The large uncertainty and the wide range of FIE values result in large uncertainties in aerosol indirect radiative forcing estimates (McComiskey and Feingold, 2008). Narrowing uncertainties in measures of aerosol–cloud interactions (ACIs) and developing well-constrained parameterizations for models requires analyses of ACIs over different climatic and aerosol regions of the Earth.

Large-scale thermodynamic conditions, such as lower tropospheric stability (LTS), significantly influence cloud development. Changes in ACIs due to different LTS have been widely investigated using observations made from the surface and from satellite remote sensing (Matsui et al., 2004; Su et al., 2010; Liu et al., 2016). However, to what degree the dependence of cloud development to aerosol perturbations is related to large-scale dynamic thermodynamic conditions is not well known. Moreover, the mechanism behind the aerosol FIE is that aerosols affect the cloud droplet number and the cloud DER through their role as CCN, which is determined by the aerosol particle size, number concentration, chemical composition, amount of water vapor, and meteorology (Sekiguchi et al., 2003; Wang et al., 2008; Qiu et al., 2017; Yang et al., 2019). Liu and Li (2018b) reported a significant influence of aerosol hygroscopicity on the magnitude of the aerosol FIE when aerosol optical quantities are used to estimate the FIE. The role of aerosol size and num-

ber concentrations on the FIE has also been examined (Garrett et al., 2004; Komppula et al., 2005; Anttila et al., 2009). Garrett et al. (2004) indicated a weak sensitivity of FIE to aerosols with small particle sizes but a stronger sensitivity to aerosols with relatively large sizes. However, the question of how sensitive cloud properties are to aerosol composition in addition to aerosol loading is still under investigation (Hao et al., 2013; Portin et al., 2014).

The Department of Energy's Atmospheric Radiation Measurement (ARM) mobile facility was stationed at Cape Cod, Massachusetts, from July 2012 to June 2013 for the Two-Column Aerosol Project (TCAP) field campaign (Berg et al., 2016). Measurements of aerosol, radiation, and cloud characteristics were made at the site, which is subject to both clear and cloudy conditions as well as clean and polluted conditions. Continental, marine, and continental–marine mixed air masses commonly pass over the site. This study uses data collected during the TCAP field campaign to investigate aerosol physical, optical, and chemical properties and their influence on the dependence of cloud development on large-scale thermodynamic conditions under different air-mass conditions. Also investigated is the influence of aerosol loading on cloud properties under different air-mass conditions and the magnitude of the FIE as well as the sensitivity of the FIE to different aerosol compositions and aerosol loadings. Section 2 describes the data and methods used in this study. Section 3 presents the seasonal variations in aerosol physical, optical, and chemical properties and their influence on low warm clouds. Section 4 gives conclusions.

2 Data and methods

2.1 Aerosol properties

2.1.1 Surface aerosol properties

The optical properties of surface aerosols were measured by a suite of instruments making up the Aerosol Observing System (AOS), which is the primary ARM platform for in situ aerosol observations. The TSI-3010 condensation particle counter was used to obtain the total number concentration of condensation particles (N_a) with diameters larger than 10 nm and smaller than 3 μm . A TSI-3653 nephelometer and a Radiance Research particle soot absorption photometer (PSAP) measured the scattering (σ_s) coefficients at three wavelengths (450, 550, and 700 nm) and the absorption (σ_a) coefficients at three wavelengths (470, 528, and 660 nm), respectively, of total ($\leq 10 \mu\text{m}$) and fine-mode ($\leq 1 \mu\text{m}$) aerosol particles (Jefferson, 2011). Nephelometer and PSAP measurements have undergone calibration and quality control using the methods developed by Anderson and Ogren (1998) and Anderson et al. (1999), respectively. Measurements of σ_a at 470 nm were normalized to 450 nm to match σ_s measurements. The single-scattering albedo (SSA) of surface aerosol

particles is then calculated as $\sigma_s/(\sigma_s + \sigma_a)$ using σ_s and σ_a at 450 nm. The time resolution of the N_a , σ_s , and σ_a measurements is 1 min.

The aerosol size distribution ranging from 15 to 450 nm was measured by a scanning mobility particle sizer (SMPS) with 5 min averaging. The SMPS contains a cylindrical differential mobility analyzer (TSI Inc., model 3081) and a condensation particle counter (TSI Inc., model 3010) and is calibrated using polystyrene latex standards (Wang et al., 2003). An aerosol chemical speciation monitor (ACSM) measured the bulk chemical composition of the nonrefractory components of submicron (aerodynamic diameter ranging from ~ 40 to 1000 nm) aerosol particles (organics, sulfate, nitrate, ammonium, and chloride). The ACSM is a thermal vaporization electron impact ionization mass spectrometer built upon the same technology as the widely used aerosol mass spectrometer. Under ambient conditions, the detection limit of the mass concentration of particles is less than $0.2 \mu\text{g m}^{-3}$ for 30 min signal averaging. The ACSM is calibrated with ammonium nitrate following the method of Ng et al. (2011).

2.1.2 Columnar aerosol properties

Columnar aerosol optical depths (AODs) and Ångström exponents (AEs) were obtained from the Aerosol Robotic Network (AERONET) database (Holben et al., 1998). AODs are retrieved from direct Sun measurements with an uncertainty of 0.01–0.02 (Eck et al., 1999). This study uses Level 2.0 quality-assured and cloud-screened data.

2.2 Cloud properties

2.2.1 Cloud boundaries

Cloud-base and cloud-top heights were identified using a combination of observations from the 95 GHz W-band ARM cloud radar (WACR), the micropulse lidar (MPL), and the ceilometer (Kollias et al., 2007). The algorithm used in the cloud boundary retrieval is similar to the method developed by Clothiaux et al. (2000) based on 35 GHz millimeter cloud radar observations. Cloud and precipitation masks are determined from the WACR based on the signal-to-noise ratio thresholds determined for each time profile. An MPL cloud mask is combined with ceilometer cloud-base estimates to produce a best-estimate cloud base for each time point. The MPL and WACR cloud masks are then merged with an additional filter to remove insect returns in the lower troposphere. Insects are identified using a combination of the WACR linear depolarization ratio and reflectivity measurements. The temporal and vertical resolutions of the cloud boundary product are 5 s and 42.856 m, respectively. Cloud-base and cloud-top heights were temporally averaged to generate data at a 1 min time resolution. The cloud-base and cloud-top height uncertainties are ~ 7.5 and ~ 45 m, respectively (Zhao et al., 2012a; Garrett and Zhao, 2013).

2.2.2 Cloud microphysical properties

A two-channel narrow-field-of-view (NFOV) radiometer and a microwave radiometer profiler (MWRP) generated cloud optical depth (COD) and LWP retrievals. The cloud droplet effective radius (r_e) was calculated using the following equation:

$$\tau = \frac{3\text{LWP}}{2\rho_w r_e},$$

where ρ_w is the density of liquid water and τ is the COD in the visible. The NFOV radiometer with a 5.7° field of view measuring downwelling zenith radiances at 673 and 870 nm at a 1 s time resolution is used to retrieve COD using the method described by Chiu et al. (2010) and Liu et al. (2013). Simultaneous highly accurate AERONET sun-photometer-measured radiances (Holben et al., 1998) quantified the biases in the NFOV radiance measurements (Fig. 1). AERONET and NFOV radiances agree well at 673 and 870 nm (coefficient of correlation, r , equal to 0.99 in both cases). However, NFOV-measured zenith radiances at 673 nm are underestimated by $\sim 15\%$. Consequently, NFOV measurements at 673 nm were adjusted using the following formula:

$$F_{673,\text{adj}} = 1.1519 \cdot F_{673,\text{obs}} + 0.0007,$$

where $F_{673,\text{obs}}$ represents measured zenith radiances and $F_{673,\text{adj}}$ represents adjusted radiances at 673 nm. The total uncertainty in COD retrievals using this method is $\sim 17\%$ (Chiu et al., 2010). The COD retrievals were averaged to generate data at a 1 min resolution for matching the time resolution of the LWP retrievals. The MWRP built by the Radiometrics Corporation measures atmospheric brightness temperatures at 12 frequencies. LWPs were retrieved using brightness temperatures measured at the five K-band channels (22.235, 23.035, 23.835, 26.235, and 30.0 GHz) with a 1 min time resolution based on the statistical retrieval algorithm developed by Liljegren and Lesht (2004). Typical uncertainties in LWP retrievals from microwave radiometers are $\sim 20 \text{ g m}^{-2}$ for $\text{LWP} < 200 \text{ g m}^{-2}$ and $\sim 10\%$ for $\text{LWP} > 200 \text{ g m}^{-2}$ (Liljegren and Lesht, 2004; Dong et al., 2008).

This study considers only nonprecipitating, low warm clouds with cloud-top heights less than 3 km. LWP observations less than 40 g m^{-2} and greater than 300 g m^{-2} were excluded to avoid very thin clouds, broken cloud cover, post-precipitation conditions (McComiskey et al., 2009), and potential precipitation contamination (Dong et al., 2008).

2.3 Surface and large-scale meteorological parameters

The ARM surface meteorological system measured surface meteorological parameters during the campaign period at a 1 min resolution. The large-scale vertical motion (ω) at

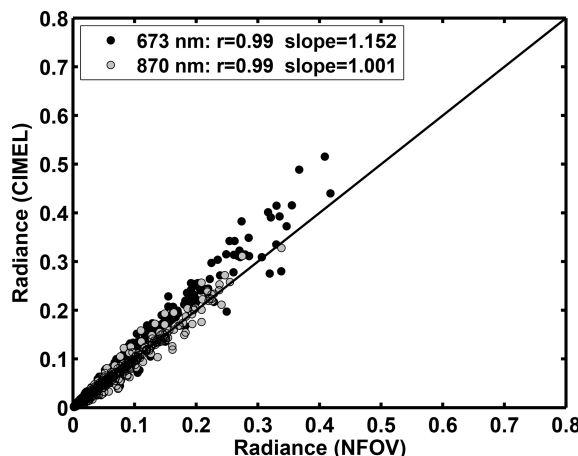


Figure 1. CIMEL sun-photometer-measured radiance as a function of narrow-field-of-view (NFOV) radiometer-measured radiance at 673 nm (black dots) and 870 nm (gray dots). The diagonal line represents the 1 : 1 line. Units are watts per steradian per squared meter ($\text{W sr}^{-1} \text{ m}^{-2}$). The legend gives the coefficient of correlation (r) and the slope of the best-fit linear regression line through each dataset.

700 hPa and LTS are used in this study to constrain large-scale dynamic and thermodynamic conditions (Su et al., 2010; Medeiros and Stevens, 2011; Liu et al., 2016). The difference between the potential temperature of the free troposphere (700 hPa) and the surface defines LTS.

To investigate the influence of aerosols on cloud properties, aerosol properties (N_a , σ_s , composition), cloud properties (COD, LWP, DER, boundary-layer height), surface meteorological parameters, and ECMWF simulations (LTS, large-scale vertical velocity) were matched according to observation time and averaged and interpolated over 1 min time intervals.

2.4 Air-mass trajectory classification

Two-day air-mass back trajectories arriving at the site at 500 m at midnight were simulated using the Hybrid Single Particle Lagrangian Integrated Trajectory model (Stein et al., 2015; Rolph, 2016). All simulated trajectories are classified into three clusters. Cluster I represents continental air masses generally originating from the continental area located to the west of the site and moving over the site. Air masses originating from the ocean area to the east of the site and directly moving over the site are the marine air masses (cluster II). Cluster III represents an air mass that has passed over both continental regions and the ocean to the site, influenced by anthropogenic and marine aerosols. During the study period, the occurrence frequencies of cluster I, II, and III air masses were 62.5 %, 15.9 %, and 21.6 %, respectively.

3 Results

3.1 Variations in aerosol properties

3.1.1 Seasonal variations in aerosol optical properties and number concentration

Figure 2 shows monthly statistics describing surface-measured σ_s for total (σ_{10}) and fine-mode (σ_1) aerosol particles and N_a . Table 1 summarizes their seasonal and annual mean values. Maxima in σ_1 and σ_{10} are found in the summer months, and minima in σ_1 and σ_{10} are found in the winter months. Fine particles dominate aerosol scattering in the summertime and are responsible for $\sim 75\%$ of the total particle scattering. The contribution of fine particle scattering to total particle scattering in other seasons ranges from $\sim 46\%$ to $\sim 54\%$, indicating that particles with diameters $\leq 1\text{ }\mu\text{m}$ and ranging from 1 to $10\text{ }\mu\text{m}$ play a similar role in aerosol scattering extinction. Monthly and seasonal variations in N_a show that maximum and minimum seasonally mean N_a values occur in spring and autumn, respectively, inconsistent with the variations in aerosol scattering coefficient. This inconsistency is probably due to seasonal differences in aerosol particle size distribution and chemical composition since aerosol extinction properties depend strongly on particle size and chemical composition. The total particle SSA shows a slight seasonal variation, suggesting smaller changes in aerosol particle absorption properties. Figure 3 shows monthly statistics describing columnar AOD and AE. Table 1 summarizes their seasonal and annual mean values. The variations in AOD and AE are consistent with the variations in surface-measured σ_s and the ratio σ_1/σ_{10} , indicating that surface aerosol properties can represent columnar aerosol properties very well at this site. Figure 4 shows monthly mean wind speeds and wind directions during the campaign period. Monthly mean wind speeds ranged from ~ 3.8 to 6.6 m s^{-1} , and southwesterly winds dominated throughout the whole year over the area. Months in summer and winter with the strongest mean surface wind speeds (e.g., June and January/February, respectively) are generally times when the contribution of fine particles to the total scattering extinction is small.

3.1.2 Aerosol optical properties under different air-mass conditions

Table 2 gives the discrepancies in aerosol properties when different air masses are in place over the site. The mean value of σ_1 is the largest/smallest under continental/marine air-mass conditions. However, σ_{10} is the largest under cluster III conditions and has similar values under cluster I and II conditions. The inconsistent variations in σ_1 and σ_{10} under different air-mass conditions can be explained by dominant particle size as indicated by the ratio σ_1/σ_{10} . When the continental air mass influenced the site, fine particles dominated

Table 1. Seasonally averaged aerosol properties during the campaign period.

	Spring	Summer	Autumn	Winter	Annual
σ_1 (Mm^{-1})	14.2 ± 14.1	33.7 ± 28.0	14.4 ± 13.6	12.8 ± 11.7	18.1 ± 19.3
σ_{10} (Mm^{-1})	31.2 ± 25.3	45.0 ± 32.9	26.5 ± 20.4	26.3 ± 23.6	31.7 ± 26.7
σ_1/σ_{10}	0.455	0.749	0.543	0.487	0.568
N_a (m^{-3})	2868 ± 2367	2498 ± 1536	2280 ± 1854	2611 ± 2108	2559 ± 2014
SSA	0.95 ± 0.04	0.96 ± 0.03	0.95 ± 0.04	0.94 ± 0.04	0.95 ± 0.04
AOD ₄₄₀	0.11 ± 0.08	0.19 ± 0.14	0.11 ± 0.11	0.08 ± 0.05	0.13 ± 0.1
AE	1.27 ± 0.40	1.65 ± 0.31	1.51 ± 0.36	1.35 ± 0.45	1.44 ± 0.40

σ_1 : scattering coefficient, fine-mode particles; σ_{10} : scattering coefficient, total; N_a : aerosol number concentration; SSA: single-scattering albedo; AOD₄₄₀: aerosol optical depth at 440 nm; AE: Ångström exponent.

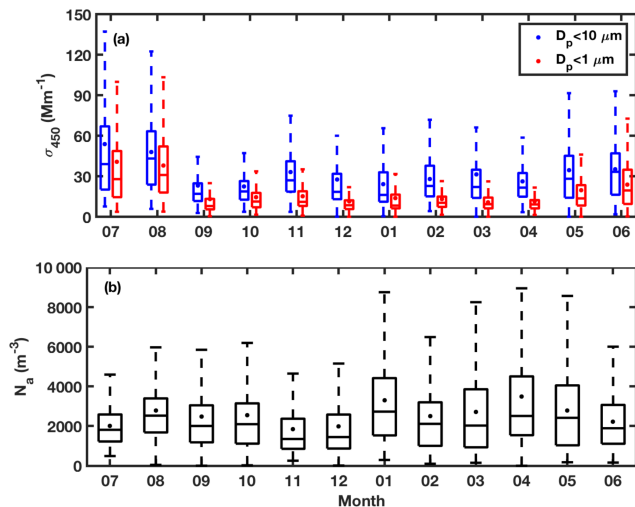


Figure 2. Monthly variations in (a) aerosol scattering coefficient at 450 nm (σ_{450}) for total (in blue, particle diameter, D_p , less than $10 \mu\text{m}$) and fine-mode (in red, D_p less than $1 \mu\text{m}$) aerosol particles and (b) aerosol particle number concentration (N_a). Box-and-whisker plots include median values (horizontal lines inside boxes), 25th and 75th percentiles (ends of boxes), 5th and 95th percentiles (ends of whiskers), and mean values (black dots). Months from left to right start at July 2012 and end at June 2013.

aerosol scattering and were responsible for $\sim 65\%$ of the total particle scattering, indicating that more anthropogenic aerosols with small particle sizes were transported to the site from continental regions to the west. The values of σ_1/σ_{10} under cluster II and III air-mass conditions show that both fine-mode and coarse-mode particles played similar roles on the total particle scattering. The variation in N_a is consistent with that in σ_1 with the largest and smallest values under cluster I and II conditions, respectively. Smaller SSA values are found under continental air-mass conditions, suggesting that more absorbing aerosols were present in this air mass than in other air masses due to the anthropogenic influence. AOD values in each air mass are similar, and the variation in AE is consistent with the variation in the ratio σ_1/σ_{10} .

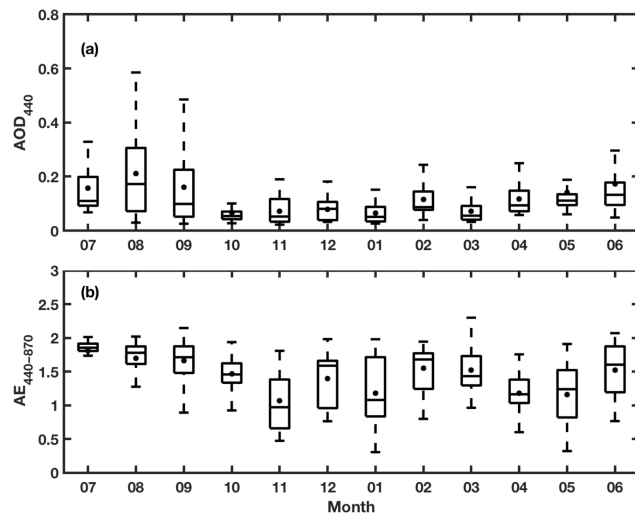


Figure 3. Monthly variations in columnar (a) aerosol optical depth at 440 nm (AOD₄₄₀) and (b) Ångström exponent (AE). Box-and-whisker plots include median values (horizontal lines inside boxes), 25th and 75th percentiles (ends of boxes), 5th and 95th percentiles (ends of whiskers), and mean values (black dots). Months from left to right start at July 2012 and end at June 2013.

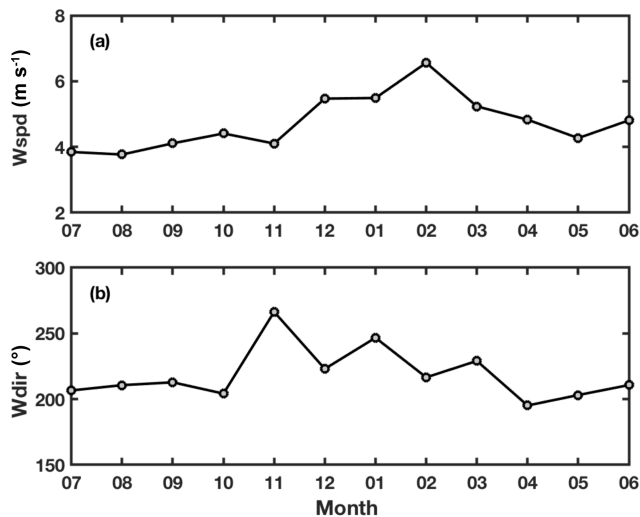
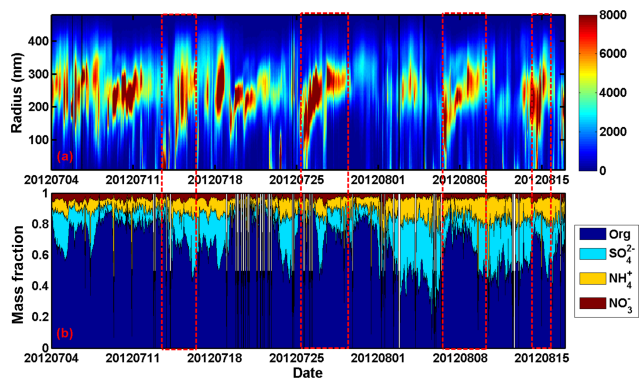
3.1.3 Aerosol chemical composition and size distribution

Figure 5 shows the size distribution and the corresponding mass fraction of organics, sulfate, ammonium, and nitrate of surface aerosol particles sampled in July and August 2012. New-particle-formation and growth periods were detected and are outlined by red rectangles in the figure. During the measurement period, fine particles containing more organics were dominant with a mean particle radius of $91.4 \pm 20.6 \text{ nm}$ and a mean organic mass fraction of 0.67 ± 0.16 . Mean mass fractions of sulfate, ammonium, and nitrate are 0.18 ± 0.11 , 0.10 ± 0.09 , and 0.04 ± 0.02 , respectively. At the beginning of new-particle-formation and growth periods, organics contributed the most to small particle sizes. Their contribution decreased as the growth period progressed to be replaced by contributions from inorganics, in particular sulfate. This is

Table 2. Means and standard deviations of aerosol and cloud properties for each air-mass cluster.

Air mass	σ_1 (Mm^{-1})	σ_{10} (Mm^{-1})	σ_1/σ_{10}	N_a (m^{-3})	SSA	AOD ₄₄₀	AE	COD	LWP (g m^{-2})	DER (μm)
I	19.8 ± 21.4	30.6 ± 26.8	0.65	2969 ± 2183	0.94 ± 0.04	0.13 ± 0.12	1.6 ± 0.4	25.7 ± 14.5	127 ± 99	7.9 ± 4.8
II	14.5 ± 14.6	30.8 ± 25.6	0.47	1788 ± 1322	0.96 ± 0.04	0.11 ± 0.08	1.3 ± 0.5	33.0 ± 18.3	243 ± 197	10.9 ± 6.6
III	16.4 ± 15.6	34.5 ± 27.2	0.48	1937 ± 1558	0.96 ± 0.03	0.12 ± 0.06	1.3 ± 0.4	26.4 ± 16.5	162 ± 121	9.8 ± 4.9

σ_1 : scattering coefficient, fine-mode particles; σ_{10} : scattering coefficient, total; N_a : aerosol number concentration; SSA: single-scattering albedo; AOD₄₄₀: aerosol optical depth at 440 nm; AE: Ångström exponent; COD: cloud optical depth; LWP: liquid water path; DER: cloud droplet effective radius.

**Figure 4.** Monthly mean (a) wind speed (Wspd) and (b) wind direction (Wdir) during the campaign period. Months from left to right start at July 2012 and end at June 2013.**Figure 5.** Time series of (a) particle size distribution and (b) mass fractions of organics (org, dark blue), sulfate (SO_4^{2-} , cyan), ammonium (NH_4^+ , yellow), and nitrate (NO_3^- , red) in aerosols sampled during July and August of 2012. Dashed red rectangles outline periods of new particle formation and growth.

possible because sulfate ions are formed during nucleation involving neutral gaseous species like ammonia and sulfuric acid (Crilley et al., 2014). Small aerosol particles generally contribute more organics to the total aerosol mass over the study site, which can also be seen in the relation between mean aerosol particle radii and organic mass fraction (Fig. 6). The strong decrease in aerosol particle size with increasing organic mass fraction has also been reported by others (Broekhuizen et al., 2006; McFiggans et al., 2006).

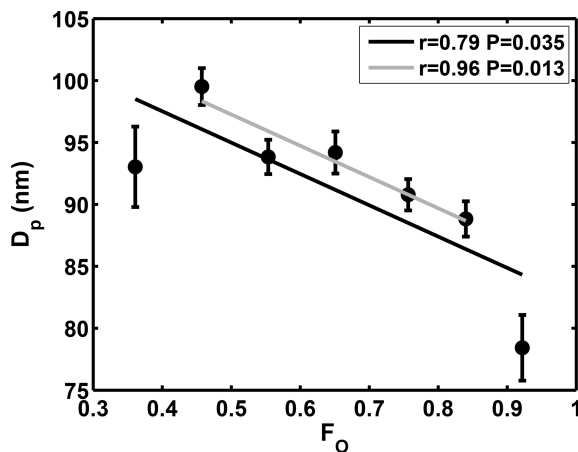


Figure 6. Mean aerosol particle radius (D_p) as a function of organic mass fraction (F_O). The black line is the linear regression line through all F_O bins. The gray line is the linear regression line for F_O bins ranging from 0.4 to 0.9, which have the most samples. Data are from July and August of 2012. The legend gives the coefficient of correlation (r) and the statistical probability (P).

3.2 Aerosol, cloud, and meteorological conditions

3.2.1 Aerosol effects on the dependence of cloud properties on meteorological conditions

Low-warm-cloud properties are sensitive to changes in thermodynamic conditions (Su et al., 2010; Medeiros and Stevens, 2011; Liu et al., 2016). Figure 7 shows cloud properties (LWP and DER) as functions of LTS under low- and high-scattering-aerosol-index (AI) conditions for continental and marine air masses. The scattering AI here is used as a proxy for CCN (Liu and Li, 2014; Sena et al., 2016) and is defined as the product of surface-measured aerosol scattering coefficients and surface-measured scattering Ångström exponents. Low and high scattering AIs are defined as the lowest and highest quarter of all scattering AI samples, respectively. The cloud properties were averaged over each 6 K LTS bin from 0 to 30 K under low- and high-scattering-AI conditions. The differences in meteorological parameters (e.g., temperature, wind speed, and relative humidity) at the surface and at 850 hPa, and in large-scale dynamic (ω) and thermodynamic parameters (LTS), are not significant under low- and high-scattering-AI conditions (figure not shown). Table 2 summarizes the means and standard deviations of cloud properties under each air-mass condition. Clouds influenced by marine air-mass conditions (cluster II) have the largest COD, LWP, and DER (33.0 ± 18.3 , $243 \pm 197 \text{ g m}^{-2}$, and $10.9 \pm 6.6 \mu\text{m}$, respectively), and clouds associated with the air mass from continental areas (cluster I) have the smallest cloud properties ($\text{COD} = 25.7 \pm 14.5$, $\text{LWP} = 127 \pm 99 \text{ g m}^{-2}$, and $\text{DER} = 7.9 \pm 4.8 \mu\text{m}$). The top panels of Fig. 7 show that LWP significantly increases with increasing LTS under low-aerosol conditions, consistent with

results from studies using surface-based measurements (e.g., Liu et al., 2016), satellite measurements (e.g., Su et al., 2010), aircraft measurements (e.g., Cecchini et al., 2016), and model simulations (e.g., Johnson et al., 2004; West et al., 2014). Johnson et al. (2004) showed that an increase in stability induces increases in the buoyancy of free-tropospheric air above the temperature inversion capping the boundary later, inhibiting the entrainment of dry air through the cloud top and increasing LWP as a result. Under high-aerosol conditions, LWP changes little as LTS increases. A likely reason is inhibited cloud droplet sedimentation due to the reduced cloud droplet size, enhancing evaporation and entrainment at the cloud top and reducing LWP (Kaufman et al., 2005; Hill et al., 2009; Zhao and Garrett, 2015; Liu et al., 2016). There are similar variations in DER with increasing LTS under low- and high-aerosol conditions (bottom panels of Fig. 7), i.e., increasing DER as LTS increases under less polluted conditions and almost constant DER as LTS increases under more polluted conditions. The changes in DER with LTS possibly reflect the changes in LWP with LTS due to the high positive correlation between LWP and DER (Zhang et al., 2011; Sporre et al., 2014). The enhanced LWP under highly stable conditions can supply the water needed for cloud droplet growth (Su et al. 2010; Zhang et al., 2011). The increase in LWP is also commonly accompanied by an increase in droplet collision–coalescence, resulting in a decrease in cloud number concentration, thus leading to an increase in DER (Kim et al., 2008; McComiskey et al., 2009; Liu et al., 2016; Zhao et al., 2018, 2019; Yang et al., 2019). Differences in LWP and DER between low- and high-LTS conditions are larger under low-pollution conditions than under high-pollution conditions. This suggests that high-aerosol concentrations can significantly weaken the thermodynamic influence on the increase in LWP and DER due to the aerosol perturbation. These results imply that the development of clouds in a highly polluted environment is inhibited even though the thermodynamic conditions may be the same as those in a much less polluted environment. The chances of precipitation are thus reduced because the rainfall frequency of warm, low clouds and LWP are highly correlated (Chen et al., 2011; Liu et al., 2013).

For all LTS bins, clouds under high-aerosol conditions have lower values of LWP and DER than clouds under low-aerosol conditions. The reduction in LWP and DER is greater in stable environments than in unstable environments, suggesting that clouds in stable environments are more affected by the aerosol perturbation than those in more unstable regimes. Studies on marine, warm clouds based on surface measurements have also shown this (Liu et al., 2016).

3.2.2 Aerosol effects on the relationships among cloud properties

Figure 8 shows the dependence of COD and DER on LWP under low- and high-scattering-AI conditions for air-

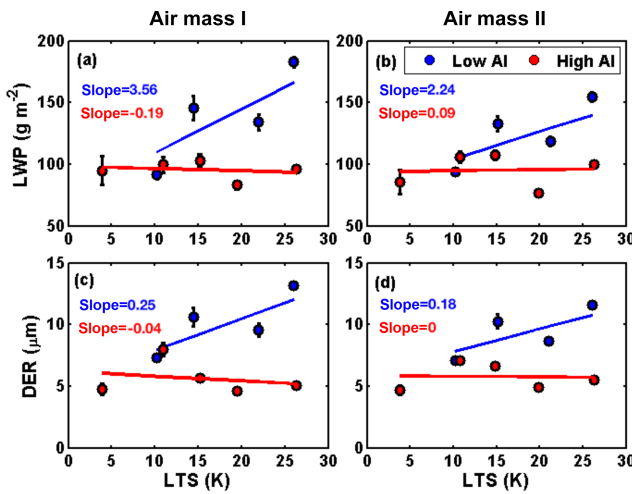


Figure 7. Liquid water path (LWP) and cloud droplet effective radius (DER) as functions of lower tropospheric stability (LTS) at low (in blue) and high (in red) scattering aerosol index (AI) levels under cluster I air-mass (**a, c**) and cluster II air-mass (**b, d**) conditions. Low and high scattering AIs are defined as the lowest and highest quarter of all scattering AI samples, respectively.

mass clusters I and II. Under high-scattering-AI conditions, COD increases sharply as LWP increases, while, under low-scattering-AI conditions, COD changes little as LWP increases due to the decrease in DER influenced by the aerosol perturbation (Fig. 8a and b). Figure 8c and d suggest that the DER is sensitive to LWP. An increase in LWP leads to a significant increase in the size of cloud droplets (Zhang et al., 2011; Sporre et al., 2014). The increase in DER with LWP is more rapid under low-scattering-AI conditions than under high-scattering-AI conditions. This is because there is a limit to the size a cloud droplet can reach when a given amount of water is shared among a large number of particles (Zhang et al., 2011). High-aerosol-loading conditions weaken the increase in DER and strengthen the increase in COD as LWP increases, indicating that aerosols have an impact on the COD–LWP and DER–LWP relationships.

Figure 8 also shows that, across all LWP bins, COD is larger and DER is smaller under high-scattering-AI conditions than under low-scattering-AI conditions, consistent with the Twomey effect. The large differences between COD under low- and high-scattering-AI conditions at high LWP values (Fig. 8a and b) and between DER under low- and high-scattering-AI conditions at high LWP values (Fig. 8c and d) suggest that when clouds have large LWPs, aerosols further inhibit the growth of cloud droplets. Under high-aerosol-loading conditions, more aerosol particles are activated into CCN, resulting in a rapid increase in cloud droplet concentration as LWP increases. However, under low-aerosol-loading conditions, cloud droplet concentrations increase slowly as LWP increases due to the lack of a CCN source, so the size

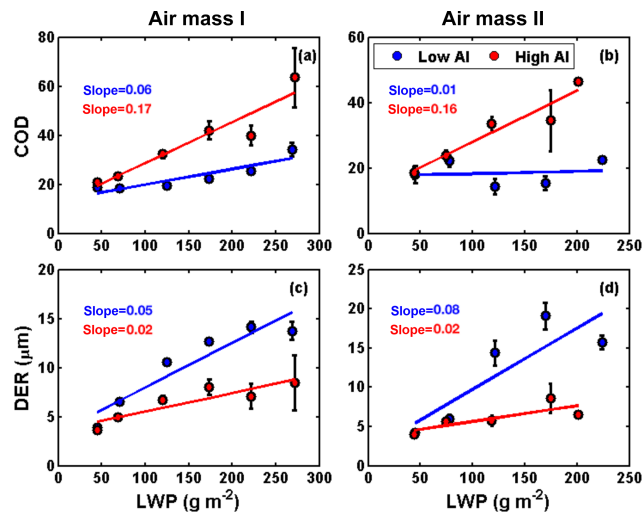


Figure 8. Cloud optical depth (COD) and cloud droplet effective radius (DER) as a function of liquid water path (LWP) at low (in blue) and high (in red) scattering aerosol index (AI) levels under cluster I air-mass (**a, c**) and cluster II air-mass (**b, d**) conditions. Low and high scattering AIs are defined as the lowest and highest quarter of all scattering AI samples, respectively.

of cloud droplets increases rapidly as LWP increases (Zhang et al., 2011).

3.3 Aerosol effect on cloud properties

3.3.1 Variations in cloud properties with aerosol loading under different air-mass conditions

Figure 9 shows the probability distribution functions (PDFs) of COD, LWP, and DER under low- and high-scattering-AI conditions for air mass clusters I and II. Numbers written in each panel are the mean percentage differences in each cloud property defined as $(M_{ch} - M_{cl})/M_{cl} \times 100\%$, where M_c represents the mean value of a cloud property and subscripts h and l represent high and low scattering AI levels, respectively. The PDFs of COD, LWP, and COD under high- and low-scattering-AI conditions differ significantly for both air masses. Although the peak values of COD under low- and high-aerosol-loading conditions are similar, clouds under more polluted conditions have more large values of COD than those under less polluted conditions. There are 24.2 % and 21.9 % increases in COD for cluster I and II respectively. For the low-aerosol-loading case, the PDF of LWP shows a broad maximum with values from 50 to 180 g m^{-2} (Fig. 9c) and 80 to 230 g m^{-2} (Fig. 9d) for cluster I and II air masses, respectively. The high-aerosol-loading cases, conversely, have narrower PDFs with distinct peaks between 60 and 70 g m^{-2} . Under high-scattering-AI conditions, the LWP decreases on the order of 30 % and 45 % from the values under low-scattering-AI conditions for the cluster I and II air masses, respectively. Under both air-mass conditions,

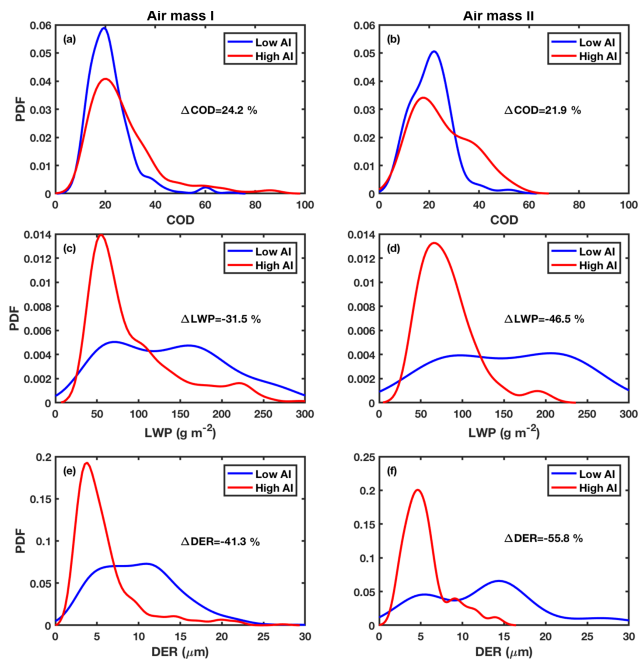


Figure 9. From top to bottom, probability distribution functions (PDFs) of cloud optical depth (COD), liquid water path (LWP), and cloud droplet effective radius (DER) at low (L, in blue) and high (H, in red) scattering aerosol index (AI) levels for the cluster I air mass (**a, c, e**) and the cluster II air mass (**b, d, f**). The mean percentage difference in COD, LWP, and DER between high and low scattering AI levels (calculated as $[(H-L)/L] \times 100\%$) is given in each panel.

there is a sharp shift in DER toward smaller values when going from high-aerosol-loading conditions to low-aerosol-loading conditions. Under low-aerosol-loading conditions, the DER values show a broad range with generally higher values varying between 5 and 12 μm for the cluster I air mass and peaking around 15 μm for the cluster II air mass. Under high-aerosol-loading conditions, the PDF of DER for both air masses is significantly narrower, with most of the values less than 10 μm and with peak values around 5 μm . From low- to high-aerosol-loading conditions, DER decreases by $\sim 40\%$ (for cluster I) and $\sim 55\%$ (for cluster II). As mentioned before, whether low- or high-scattering-AI conditions are in place, meteorological parameters and large-scale dynamic and thermodynamic parameters show little difference, suggesting that aerosols are responsible for the changes in cloud properties. In general, clouds in a marine air mass have slightly larger decreases in LWP and DER from low to high aerosol loading than those in a continental air mass.

3.3.2 Aerosol first indirect effect

The aerosol FIE is generally quantified as

$$\text{FIE} = -\frac{d\ln(\text{DER})}{d\ln(\alpha)}|_{\text{LWP}},$$

where α represents CCN or CCN proxies. The FIE represents the relative change in mean cloud DER with respect to a relative change in aerosol loading for clouds having the same LWP (Feingold et al., 2003). In some studies, the scattering AI is used as the CCN proxy (Liu and Li, 2014). Cloud samples were categorized according to their LWP values. The LWP bins range from 40 to 200 g m^{-2} in increments of 20 g m^{-2} . The choice of a small increment ensures that the LWP constraint is met in each bin. Since there were not enough samples under cluster II air-mass conditions, only FIE for clouds and aerosols under cluster I air-mass conditions are calculated. Only those cases with sample numbers greater than 50 per bin where the calculated values of FIE are statistically significant at the 95 % confidence level ($P = 0.05$) are analyzed here. Figure 10a shows DER as function of scattering AI for clouds with LWP ranging from 120 to 140 g m^{-2} . It illustrates how the FIE is estimated. There is a significant decrease in DER as the scattering AI increases. For this case, the magnitude of the FIE is 0.26 with an uncertainty of 0.09. Figure 10b shows the magnitudes and uncertainties of FIE calculated in each LWP bin. Numbers above each bar are the number of samples that went into the calculation of the FIE in each LWP bin. The magnitude of the FIE changes from 0.07 ± 0.03 to 0.26 ± 0.09 with the smallest value found in the LWP bin of 40–60 g m^{-2} and the largest value found in the LWP bin of 120–140 g m^{-2} . The mean value of FIE during the study period based on all LWP bins is 0.16 ± 0.06 . The values of FIE in each LWP bin increase with increasing LWP, especially for LWP less than 140 g m^{-2} . This is consistent with results from previous studies (e.g., Pandithurai et al., 2009; Sporre et al., 2014; Harikishan et al., 2016). Enhanced aerosol activation due to the increase in LWP may explain this (Zhao et al., 2012b; Painemal and Zuidema, 2013). At higher LWP values, with the availability of more CCN, more droplets can activate. The droplet number increases, but their size decreases at fixed LWP levels (Harikishan et al., 2016). Estimates of the FIE reported from all available platforms range widely and are sensitive to the definition of the aerosol burden (Lihavainen et al., 2010; Zhao et al., 2012b), the methods for retrieving cloud properties (McComiskey et al., 2009), and meteorological conditions such as vertical velocity and atmospheric stability (Feingold et al., 2003; Matsui et al., 2004; McComiskey et al., 2009; Liu et al., 2016; Yang et al., 2019). Theoretical values of the FIE lie between 0 and 0.33 (McComiskey and Feingold, 2008) with most values falling between 0.05 and 0.25 (Zhao et al., 2012b). Based on surface retrievals, Feingold et al. (2003) derived FIE values of 0.02–0.16 with a mean value of 0.10 ± 0.05 for a set of seven cases. In a study using 3 years of data from the United States Southern Great Plains, Kim et al. (2008) found that FIE values ranged from 0.04 to 0.17 over five LWP bins with a mean value of 0.09 ± 0.05 . Other estimates of FIE based on surface retrievals have been reported, e.g., 0.07 ± 0.01 for warm, marine boundary-layer clouds over the Azores (Liu et al.,

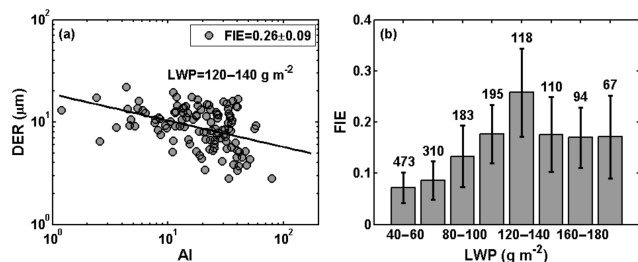


Figure 10. (a) Cloud droplet effective radius (DER) as a function of scattering aerosol index (AI) for a sample bin with a constant liquid water path (LWP) range of 120 to 140 g m^{-2} , and (b) the quantified aerosol first indirect effect (FIE) for each LWP bin. Numbers above each bar in (b) are the number of samples that went into the calculation of the FIE.

2016); 0.14 ± 0.09 for continental clouds during the monsoon period over a rural continental site in Mahabubnagar, India (Harikishan et al., 2016); and a range of 0.05 to 0.16 over the coastal region at Pt. Reyes, California (McComiskey et al., 2009). The magnitude of the FIE in this study generally falls in this range.

Examined next is the sensitivity of cloud properties to aerosol chemical composition represented by the mass fraction of organics. The aerosol number concentration is used as the CCN proxy here (Li et al., 2011; Yan et al., 2014; Liu et al., 2016) because aerosol scattering coefficient measurements were not taken during the aerosol chemical composition observation period. Three LWP bins were defined: 40–60, 60–80, and 80–100 g m^{-2} . Figure 11 shows DER as a function of N_a in each LWP bin when aerosol particle mass fractions of organics are low and high. Aerosols with low and high mass fractions of organics are defined as aerosols with mass fractions of organics less than and greater than, respectively, the mean value of the mass fraction of organics of all samples in each LWP bin. Mean values of ω and LTS in each aerosol particle mass fraction of organics category are given in the figure. Differences in ω and LTS between low and high mass fractions of organics are not significant in any LWP bin. FIE estimates when aerosol samples with low mass fractions of organics dominate are 0.10 ± 0.05 , 0.15 ± 0.06 , and 0.23 ± 0.12 (Fig. 11a–c, respectively), which are greater than the FIE estimates when aerosol samples with high mass fractions of organics dominate (0.07 ± 0.04 , 0.12 ± 0.06 , and 0.07 ± 0.05 , respectively). Clouds are more susceptible to inorganics-dominant aerosol than to organic aerosols, resulting in a greater climate forcing. The mechanism behind the aerosol indirect effect is characterized by the ability of aerosol particles to act as CCN, which is primarily governed by particle size and chemical composition (McFiggans et al., 2006). The cloud-nucleating ability of aerosol particles is significantly greater when the aerosol particles are larger and composed of more inorganic compounds than when they are small and composed of more organic com-

pounds (Dusek et al., 2006; Liu et al., 2011). This study (see Fig. 5) and others have demonstrated that aerosols containing more organic particles are generally smaller than those with more inorganic particles (Broekhuizen et al., 2006; McFiggans et al., 2006; Zhang et al., 2011) and that organic particles are generally less CCN-active than inorganic particles (Raymond and Pandis, 2002; Zhang et al., 2011). This can partly explain the smaller FIE values induced by aerosols with large mass fractions of organics.

4 Conclusions

Twelve months (July 2012–June 2013) of measurements of aerosol and cloud properties, as well as meteorological variables, were acquired during the Two-Column Aerosol Project field campaign at Cape Cod, Massachusetts. The goal of this study is to characterize aerosol physical, optical, and chemical composition properties and to determine their influences on cloud properties and the dependence of cloud development on large-scale thermodynamic conditions. Also examined was the magnitude of the aerosol FIE and the sensitivities of cloud properties to aerosol composition in addition to aerosol loading.

The maximum and minimum in σ_1 and σ_{10} were found in summer and winter, respectively. Fine particles dominated aerosol scattering in the summer and contributed toward $\sim 75\%$ of the total particle scattering. In other seasons, fine particles contributed toward $\sim 45\%–54\%$ of the total particle scattering. The maximum and minimum mean values of N_a occurred in spring and autumn, which is not consistent with the variation in σ_s . The variation in AOD is consistent with the variation in surface-measured σ_s and inconsistent with the variation in N_a . This suggests that a large number of particles with less optical sensitivity were present. Months with strong mean surface wind speeds were generally associated with small σ_s and a small contribution of fine particles to the total scattering extinction but relatively large aerosol number concentrations. This suggests that strong surface winds had ushered in from the inland continental region more optically insensitive aerosols of small particle size. For all new-particle-formation and growth cases considered in this study, a large contribution of organics to small particles was observed, which then decreased during the particle growth period.

Under low-scattering-AI conditions, LWP and DER significantly increased as LTS increased, but, under high-scattering-AI conditions, LWP and DER changed little. Differences in LWP and DER between low- and high-LTS conditions were larger under light pollution than under heavy pollution. This suggests that the dependence of cloud properties is weakened by the aerosol perturbation. The reduction in LWP and DER was greater in stable environments than in unstable environments, indicating that clouds in stable environments are more influenced by aerosol pertur-

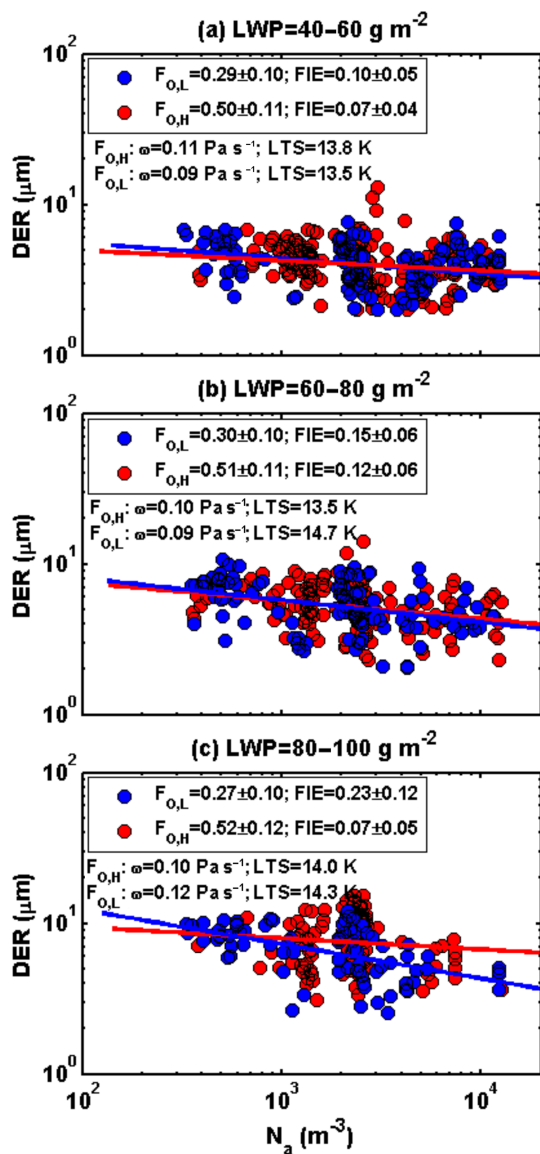


Figure 11. Cloud droplet effective radius (DER) as a function of aerosol number concentration (N_a) at low (in blue) and high (in red) levels of mass fraction of organics in three liquid water path (LWP) bins: (a) $40\text{--}60\text{ g m}^{-2}$, (b) $60\text{--}80\text{ g m}^{-2}$, and (c) $80\text{--}100\text{ g m}^{-2}$. Linear regression lines through each set of data are drawn. $F_{o,l}$ and $F_{o,h}$ are defined as the means of values less than and greater than, respectively, the mean value of the mass fraction of organics from all samples in each LWP bin. The legend gives the mean values of $F_{o,l}$ and $F_{o,h}$ with their standard deviations and the magnitudes of the FIE with their uncertainties. Mean values of vertical velocity (ω) and lower tropospheric stability (LTS) corresponding to $F_{o,l}$ and $F_{o,h}$ levels in each LWP bin are also given.

butions than those in more unstable regimes. DER significantly increased with increasing LWP under low-aerosol conditions but slowly increased as LWP increased under high-aerosol conditions. Under high-scattering-AI conditions, COD sharply increased with increasing LWP, but, un-

der low-scattering-AI conditions, the increase was slower. This suggests that aerosols can influence the interdependence of cloud properties.

Analyses of the PDFs of COD, LWP, and DER under low- and high-aerosol-loading conditions in continental air mass (clusters I) and oceanic air mass (clusters II) suggest that high aerosol loading can increase COD and decrease LWP and DER and narrow the distributions of LWP and DER. The magnitude of FIE estimated under continental air-mass conditions ranged from 0.07 ± 0.03 to 0.26 ± 0.09 with a mean value of 0.16 ± 0.03 and showed an increasing trend as LWP increased. The magnitude of the FIE estimated for aerosols with a low fraction of organics was larger than that for aerosols with a high mass of organics. This suggests that clouds over regions dominated by inorganic aerosols are more susceptible to aerosol perturbations, resulting in larger climate forcing, than clouds over regions dominated by organic aerosols.

Data availability. Data used in this study can be accessed from the public FTP of the Atmospheric Radiation Measurement (ARM) at <https://www.archive.arm.gov/discovery/> (last access: 20 December 2018).

Author contributions. JL performed the calculations, analyzed the data, and wrote the paper. ZL contributed to the writing of paper and in scientific discussions of the study.

Competing interests. The authors declare that they have no conflict of interest.

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